blueTEV, lyophilized formulation

Cysteine protease from Tobacco Etch Virus Cat. no. P2020-137



Product Information

Protein: blueTEV, lyophilized formulation (~ 53.7 kDa)

Uniprot#: QOGDU8

Sequence: KGPRDYNPISSSICHLTNESDGHTTSLYGIGFGPFIITNKHLFRRNNGTLVVQSLHGVFK

VKDTTTLQQHLVDGRDMIIIRMPKDFPPFPQKLKFREPQREERICLVTTNFQTKSMSSMV SDTSCTFPSGDGIFWKHWIQTKDGQCGSPLVSTRDGFIVGIHSASNFTNTNNYFTSVPKN

FMELLTNQEAQQWVSGWRLNADSVLWGGHKVFMVKPEEPFQPVKEATQLMNE

Methionine at pos. 1 might be present due to cloning constraints, C-terminal His-tag and

BFP-fusion not shown in sequence.

Source: Recombinantly expressed in *E.coli*.

Tag(s): BFP, N-terminal and His-Tag, C-terminal

Purification: Purified by affinity chromatography and subsequent buffer exchange.

Formulation: 50 mM Tris, 150 mM NaCl, 0.5 mM EDTA, 5 % (w/v) Trehalose; pH 8.0.

Lyophilized, stored at -80 $^{\circ}\text{C}$ and shipped at ambient temperature.

We recommend reconstituting the enzyme in 40 % Glycerol (w/v).

In case of 1000 Units, add 50 μ l of 40 % Glycerol (w/v). In case of 10'000 Units, add 500 μ l of 40 % Glycerol (w/v).

Purity: > 85 % (will be determined by densitometry of Coomassie stained gel, example next page)

Specific Activity: ≥ 20 Units/µl (determined by cleavage of control protein)

≥ 0.25 µmol/min/mg (determined by cleavage of labeled peptide (Fluorometric assay), TEV

Protease Activity Kit (Abcam))

Unit definition: One unit of blueTEV cleaves > 85 % of 3 µg of a fusion protein in 1 hour at room

temperature. Cleavage overnight increases cleavage efficiency to > 99 %. It is

recommended to optimize cleavage conditions for each protein by varying the amount of

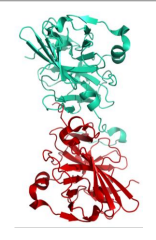
blueTEV, reaction time, or temperature.

Concentration: Will be determined by BCA-Assay.

Long-term storage: Recommended at -20 °C.

Background Information:

blueTEV represents the catalytic domain of the nuclear inclusion a (NIa) protein with a molecular weight of 27 kDa encoded by the plant virus Tobacco Etch Virus. "blue" indicates fusion of the protease to blue fluorescent protein (BFP), which leads to increased stability and solubility of TEV protease. Moreover, blueTEV has been optimized by site directed mutagenesis to prevent autocatalytic cleavage. blueTEV is a highly site-specific cysteine protease that recognizes the amino acid sequence Glu-Asn-Leu-Tyr-Phe-Gln-(Gly/Ser) [ENLYFQ(G/S)] and cleaves between the residues Gln and Gly/Ser. The most commonly used recognition sequence is ENLYFQG. In biotechnology,



Structural model of blueTEV, lyophilized formulation

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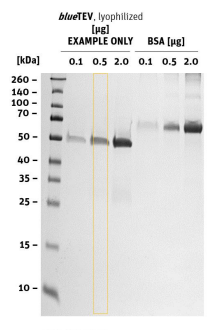


Product Information

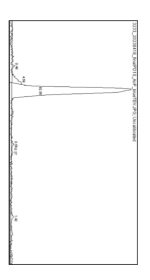
blueTEV is a versatile enzyme to remove affinity tags from recombinant proteins with high specificity and activity over a wide range of pH, ionic strength and temperatures between 4 °C and 30 °C. The optimal temperature for cleavage is 30 °C. It is recommended to improve cleavage efficiency for each fusion protein by varying the amount of recombinant blueTEV, reaction time, or incubation temperature. The great advantage of blueTEV is its facile removal after cleavage reaction by immobilized metal affinity chromatography (IMAC) since it is equipped with a Histag. Furthermore, the removal of blueTEV can be monitored instantly by detection of fluorescence in solution - this easy, fast and sensitive method omits

time-consuming SDS-PAGE or Western blot analysis. If the blue fluorescence of blueTEV is not suitable and you prefer green fluorescence as readout, check our greenTEV protease.

Quality Information (provided for each lot):



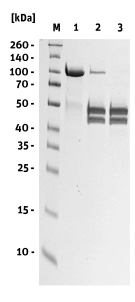




	Area	Percent
1	37.071	0.487
2	349.506	4.592
3	7045.527	92.562
4	42.778	0.562
5	28.364	0.373
6	108.435	1.425

Histogram (of marked lane in gel picture)

Test Cleavage:



P2020-131 Cleavage and tag control protein (90.5 kDa):

Start of cleavage reaction Cleavage reaction (1 h) End of cleavage reaction (24 h)

12 % SDS-PAGE & coll. Coomassie staining.

SDS-PAGE/Coll.Coomassie